Welcome to STN International! Enter x:x

## LOGINID:SSSPTA1642BJF

## PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

* *	* *	* *	* *	* Welcome to STN International * * * * * * * * *
NEW	1			Web Dans for CTN Conings Cabadala N. America
				Web Page for STN Seminar Schedule - N. America
				STN pricing information for 2008 now available
NEWS	3	JAN	16	CAS patent coverage enhanced to include exemplified
				prophetic substances
NEWS	4	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new
	_			custom IPC display formats
				MARPAT searching enhanced
NEWS	6	JAN	28	USGENE now provides USPTO sequence data within 3 days
				of publication
				TOXCENTER enhanced with reloaded MEDLINE segment
				MEDLINE and LMEDLINE reloaded with enhancements
				STN Express, Version 8.3, now available
				PCI now available as a replacement to DPCI
				IFIREF reloaded with enhancements
				IMSPRODUCT reloaded with enhancements
NEWS	13	FEB	29	WPINDEX/WPIDS/WPIX enhanced with ECLA and current
				U.S. National Patent Classification
NEWS	14	MAR	31	IFICDB, IFIPAT, and IFIUDB enhanced with new custom
				IPC display formats
NEWS	15	MAR	31	CAS REGISTRY enhanced with additional experimental
				spectra
NEWS	16	MAR	31	CA/CAplus and CASREACT patent number format for U.S.
				applications updated
				LPCI now available as a replacement to LDPCI
				EMBASE, EMBAL, and LEMBASE reloaded with enhancements
				STN AnaVist, Version 1, to be discontinued
NEWS	20	APR	15	WPIDS, WPINDEX, and WPIX enhanced with new
				predefined hit display formats
				EMBASE Controlled Term thesaurus enhanced
				IMSRESEARCH reloaded with enhancements
NEWS	23	MAY	30	INPAFAMDB now available on STN for patent family
	0.1		2.0	searching
NEWS	24	MAI	30	DGENE, PCTGEN, and USGENE enhanced with new homology
MELLO	0.5	77737	0.0	sequence search option
				EPFULL enhanced with 260,000 English abstracts
				KOREAPAT updated with 41,000 documents USPATFULL and USPAT2 updated with 11-character
MEMS	21	JUN	13	patent numbers for U.S. applications
MENTO	20	TIINI	10	CAS REGISTRY includes selected substances from
NEWS	28	JUN	19	web-based collections
	20	77777	2.5	CA/CAplus and USPAT databases updated with IPC
NEWS	29	JUN	25	reclassification data
NIDLIC	20	TIBI	20	AEROSPACE enhanced with more than 1 million U.S.
NEWS	50	OON	50	
NIDLIC	2.1	TIINI	20	patent records EMBASE, EMBAL, and LEMBASE updated with additional
NEWS	21	OUN	30	options to display authors and affiliated
				operons to display authors and allitiated
	NEWS NEWS NEWS NEWS NEWS NEWS NEWS NEWS	NEWS 2 2 3 NEWS 4 NEWS 5 NEWS 5 NEWS 10 NEWS 11 NEWS 14 NEWS 15 NEWS 16 NEWS 17 NEWS 18 NEWS 20 NEWS 20 NEWS 20 NEWS 21 NEWS 2	NEWS 2 JAN NEWS 2 JAN NEWS 4 JAN NEWS 5 JAN NEWS 6 JAN NEWS 7 JAN NEWS 8 JAN NEWS 10 FEB NEWS 10 FEB NEWS 10 FEB NEWS 12 APR NEWS 14 MAR NEWS 15 APR NEWS 16 JAPR NEWS 17 APR NEWS 18 APR NEWS 20 APR NEWS 21 APR NEWS 21 APR NEWS 22 APR NEWS 22 APR NEWS 21 APR NEWS 22 APR NEWS 22 APR NEWS 21 APR NEWS 22 APR NEWS 22 APR NEWS 21 APR NEWS 22 APR NEWS 22 APR NEWS 22 APR NEWS 22 APR NEWS 21 APR NEWS 22 APR NEWS 23 APR NEWS 24 APR NEWS 25 APR NEWS 26 APR NEWS 27 JUN NEWS 26 APR NEWS 27 JUN NEWS 27 JUN NEWS 28 APR NEWS 29 APR NEWS 29 APR NEWS 20 APR NEWS 20 APR NEWS 20 APR NEWS 21 APR NEWS 22 APR	NEWS 2 JAN 02 RINEWS 3 JAN 28 RINEWS 5 JAN 28 RINEWS 6 JAN 28 RINEWS 6 JAN 28 RINEWS 7 JAN 28 RINEWS 7 JAN 28 RINEWS 10 FEB 20 RINEWS 11 FEB 25 RINEWS 12 FEB 25 RINEWS 15 MAR 31 RINEWS 16 MAR 31 RINEWS 16 MAR 31 RINEWS 17 MAR 31 RINEWS 18 MAR 31 RINEWS 18 MAR 31 RINEWS 18 MAR 31 RINEWS 19 APR 04 RINEWS 20 APR 28 RINEWS 21 APR 28 RINEWS 22 APR 28 RINEWS 22 APR 28 RINEWS 22 APR 28 RINEWS 21 APR 28 RINEWS 21 APR 28 RINEWS 22 APR 28 RINEWS 25 APR 28 RINEWS 26 JUN 06 RINEWS 26 JUN 06 RINEWS 27 JUN 13 RINEWS 29 JUN 25 RINEWS 20 JUN 25 RINEWS 30 JUN 30 RINEWS 20 JUN 25 RINEWS 30 JUN 30 RINEWS 20 JUN 25 RINEWS 30 JUN 30 JUN 30 RINEWS 30 JUN

organizations

NEWS 32 JUN 30 STN on the Web enhanced with new STN AnaVist

Assistant and BLAST plug-in

NEWS 33 JUN 30 STN AnaVist enhanced with database content from EPFULL

NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3. AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

NEWS HOURS STN Operating Hours Plus Help Desk Availability

NEWS LOGIN Welcome Banner and News Items

NEWS IPC8 For general information regarding STN implementation of IPC 8

Enter NEWS followed by the item number or name to see news on that specific topic.

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\* \* \* \* \* \* \* \* \* \* \* \* \* \* \* \* STN Columbus \* \* \* \* \* \* \* \* \* \* \* \* \* \* \* \* \* \*

FILE 'HOME' ENTERED AT 08:59:51 ON 07 JUL 2008

=> file req

COST IN U.S. DOLLARS SINCE FILE TOTAL SESSION ENTRY FULL ESTIMATED COST 0.21 0.21

FILE 'REGISTRY' ENTERED AT 09:00:16 ON 07 JUL 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 6 JUL 2008 HIGHEST RN 1032827-24-9 DICTIONARY FILE UPDATES: 6 JUL 2008 HIGHEST RN 1032827-24-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> E "EPOTHILONE B"/CN 25

```
E1
           1 EPOTHILONE A8/CN
E2
           1
                EPOTHILONE A9/CN
E3
           1 --> EPOTHILONE B/CN
```

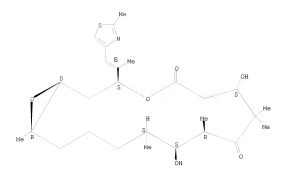
E4 1 EPOTHILONE B (12R,13R) ACETONIDE/CN

1 EPOTHILONE B A-EPOXIDE/CN E5

E6 1 EPOTHILONE B ACID/CN

```
1
                   EPOTHILONE B HYDROXYLASE/CN
E7
E8
             1
                   EPOTHILONE B HYDROXYLASE (AMYCOLATOPSIS ORIENTALIS GENE EBH)/CN
E9
             1
                   EPOTHILONE B N-OXIDE/CN
E10
                  EPOTHILONE B10/CN
             1
E11
                  EPOTHILONE C/CN
             1
           1 EPOTHILONE C BIS(TERT-BUTYLDIMETHYLSILYL) ETHER/CN
1 EPOTHILONE C/D 12,13-EPOXIDASE/CN
1 EPOTHILONE C/D MONOXYGENASE/CN
1 EPOTHILONE C/D SYNTHETASE/CN
1 EPOTHILONE C/D C/CN
1 EPOTHILONE C/CN
E12
E13
E14
E15
E16
E17
E18
            1
                  EPOTHILONE C3/CN
E19
            1
                  EPOTHILONE C4/CN
E20
            1
                   EPOTHILONE C5/CN
E21
            1
                   EPOTHILONE C6/CN
E22
            1
                   EPOTHILONE C7/CN
E23
            1
                   EPOTHILONE C8/CN
             1
E24
                   EPOTHILONE C9/CN
             1
E25
                   EPOTHILONE D/CN
=> S E3
L1
             1 "EPOTHILONE B"/CN
=> S L1 EXA SAM
SAMPLE IS IGNORED AS A SCOPE FOR THIS SEARCH
              1 "EPOTHILONE B"/CN
=> DTS L2 1 SAM
THE ESTIMATED COST FOR THIS REQUEST IS 1.04 U.S. DOLLARS
DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y) /N:Y
    ANSWER 1 OF 1 REGISTRY COPYRIGHT 2008 ACS on STN
L2
TN
     4,17-Dioxabicyclo[14.1.0]heptadecane-5,9-dione, 7,11-dihydroxy-
     8, 8, 10, 12, 16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-
     , (1S, 3S, 7S, 10R, 11S, 12S, 16R) -
MF
     C27 H41 N O6 S
```

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

=>

---Logging off of STN---

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 13.64 13.85

STN INTERNATIONAL LOGOFF AT 09:03:08 ON 07 JUL 2008

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID:SSSPTA1642BJF

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
Welcome to STN International
NEWS 1
                 Web Page for STN Seminar Schedule - N. America
NEWS 2 JAN 02
                STN pricing information for 2008 now available
NEWS 3 JAN 16 CAS patent coverage enhanced to include exemplified
                 prophetic substances
NEWS 4
        JAN 28
                USPATFULL, USPAT2, and USPATOLD enhanced with new
                 custom IPC display formats
NEWS 5
        JAN 28
                MARPAT searching enhanced
        JAN 28 USGENE now provides USPTO sequence data within 3 days
                 of publication
NEWS 7
        JAN 28
                TOXCENTER enhanced with reloaded MEDLINE segment
NEWS 8
        JAN 28 MEDLINE and LMEDLINE reloaded with enhancements
NEWS 9 FEB 08 STN Express, Version 8.3, now available
NEWS 10 FEB 20 PCI now available as a replacement to DPCI
NEWS 11 FEB 25
                IFIREF reloaded with enhancements
NEWS 12
                IMSPRODUCT reloaded with enhancements
        FEB 25
NEWS 13 FEB 29
                WPINDEX/WPIDS/WPIX enhanced with ECLA and current
                 U.S. National Patent Classification
NEWS 14 MAR 31
                 IFICDB, IFIPAT, and IFIUDB enhanced with new custom
                 IPC display formats
NEWS 15 MAR 31
                 CAS REGISTRY enhanced with additional experimental
                 spectra
NEWS 16 MAR 31
                CA/CAplus and CASREACT patent number format for U.S.
                 applications updated
NEWS 17 MAR 31
                LPCI now available as a replacement to LDPCI
NEWS 18 MAR 31
                EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 19
        APR 04
                STN AnaVist, Version 1, to be discontinued
NEWS 20
        APR 15 WPIDS, WPINDEX, and WPIX enhanced with new
                 predefined hit display formats
        APR 28 EMBASE Controlled Term thesaurus enhanced
NEWS 21
NEWS 22
        APR 28 IMSRESEARCH reloaded with enhancements
NEWS 23
        MAY 30
                INPAFAMDB now available on STN for patent family
                 searching
NEWS 24
        MAY 30
                DGENE, PCTGEN, and USGENE enhanced with new homology
                 sequence search option
NEWS 25
        JUN 06
                EPFULL enhanced with 260,000 English abstracts
NEWS 26
        JUN 06
                KOREAPAT updated with 41,000 documents
NEWS 27
        JUN 13 USPATFULL and USPAT2 updated with 11-character
                 patent numbers for U.S. applications
NEWS 28
        JUN 19
                CAS REGISTRY includes selected substances from
                 web-based collections
NEWS 29
        JUN 25 CA/CAplus and USPAT databases updated with IPC
                 reclassification data
NEWS 30
        JUN 30 AEROSPACE enhanced with more than 1 million U.S.
                 patent records
        JUN 30 EMBASE, EMBAL, and LEMBASE updated with additional
NEWS 31
                 options to display authors and affiliated
                 organizations
NEWS 32
        JUN 30
                STN on the Web enhanced with new STN AnaVist
                 Assistant and BLAST plug-in
NEWS 33
        JUN 30 STN AnaVist enhanced with database content from EPFULL
NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3.
             AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.
              STN Operating Hours Plus Help Desk Availability
NEWS HOURS
NEWS LOGIN
              Welcome Banner and News Items
NEWS IPC8
             For general information regarding STN implementation of IPC 8
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Enter NEWS followed by the item number or name to see news on that

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FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008

=> file pctfull

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION

0.21

FULL ESTIMATED COST 0.21

FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008 COPYRIGHT (C) 2008 Univentio

FILE LAST UPDATED: 4 JUL 2008 <20080704/UP>
FILE COVERS 1978 TO DATE

>>> IMAGES ARE AVAILABLE ONLINE AND FOR EMAIL-PRINTS <<<

>>> NEW FIELD UPTX, FIELD /EW NO LONGER AVAILBLE - SEE HELP CHANGE <<<

=> s epothilon?

L1 2484 EPOTHILON?

=> s 11/ab or 11/ti

144 EPOTHILON?/AB 129 EPOTHILON?/TI

L2 159 (EPOTHILON?/AB) OR (EPOTHILON?/TI)

=> s 12 not py>2001 817323 PY>2001

L3 53 L2 NOT PY>2001

=> s combination and 13

567168 COMBINATION 264042 COMBINATIONS

617900 COMBINATION

(COMBINATION OR COMBINATIONS)

L4 33 COMBINATION AND L3

=> d ibib 1-5

L4 ANSWER 1 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN

ACCESSION NUMBER: 2001092255 PCTFULL ED 20020826

TITLE (ENGLISH): EPOTHILONE DERIVATIVES AND METHODS FOR MAKING

AND USING THE SAME
TITLE (FRENCH): DERIVES D'EPOTHILONE, PROCEDES DE PRODUCTION

ET METHODES D'UTILISATION

INVENTOR(S): SANTI, Daniel;
FARDIS, Maria;

ASHLEY, Gary

PATENT ASSIGNEE(S): KOSAN BIOSCIENCES, INC.;

SANTI, Daniel; FARDIS, Maria; ASHLEY, Gary

DOCUMENT TYPE:

Patent

PATENT	INFORMATION:	

PATENT INFORMATION:	NUMBER KIND	DATE
	WO 2001092255 A2	
DESIGNATED STATES	WO 2001092255 AZ	20011206
W:		BB BG BR BY BZ CA CH CN CO C ES FI GB GD GE GH GM HR HU I
		KZ LC LK LR LS LT LU LV MA M
		PL PT RO RU SD SE SG SI SK S
		UZ VN YU ZA ZW GH GM KE LS M AZ BY KG KZ MD RU TJ TM AT B
		GR IE IT LU MC NL PT SE TR B
PRIORITY INFO.:	BJ CF CG CI CM GA GN GW US 2000-60/207,655	
	US 2000-60/218,260	20000714
APPLICATION INFO.:	US 2000-60/231,552 WO 2001-US15763 A	20010515
L4 ANSWER 2 OF 33		
ACCESSION NUMBER:		
TITLE (ENGLISH):	PRODUCTION OF POLYKETID	
TITLE (FRENCH): INVENTOR(S):	PRODUCTION DE POLYKETID ARSLANIAN, Robert, L.;	ES
	ASHLEY, Gary;	
	FRYKMAN, Scott; JULIEN, Bryan;	
	KATZ, Leonard;	
	KHOSLA, Chaitan; LAU, Janice;	
	LICARDI, Peter, J.;	
	REGENTIN, Rika; SANTI, Daniel;	
	TANG, Li	
PATENT ASSIGNEE(S):	KOSAN BIOSCIENCES, INC. ARSLANIAN, Robert, L.;	;
	ASHLEY, Gary;	
	FRYKMAN, Scott; JULIEN, Bryan;	
	KATZ, Leonard;	
	KHOSLA, Chaitan;	
	LAU, Janice; LICARDI, Peter, J.;	
	REGENTIN, Rika;	
	SANTI, Daniel; TANG, Li	
DOCUMENT TYPE:	Patent	
PATENT INFORMATION:	NUMBER KIND	DATE
	WO 2001083800 A2	20011108
DESIGNATED STATES W:	AE AG AL AM AT AU A7 BA	BB BG BR BY BZ CA CH CN CR C
	CZ DE DK DM DZ EE ES FI	GB GD GE GH GM HR HU ID IL I
		LK LR LS LT LU LV MA MD MG M RO RU SD SE SG SI SK SL TJ T
	TR TT TZ UA UG US UZ VN	YU ZA ZW GH GM KE LS MW MZ S
		KG KZ MD RU TJ TM AT BE CH C IT LU MC NL PT SE TR BF BJ C
	CG CI CM GA GN GW ML MR	NE SN TD TG
PRIORITY INFO.:	US 2000-09/560,367 US 2000-60/232,696	20000428 20000914
	US 2000-60/257,517	20001221

US 2001-09/825,856 20010403 US 2001-09/825,876 20010403 US 2001-60/269,020 20010413 WO 2001-US13793 A 20010426 APPLICATION INFO . . ANSWER 3 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN ACCESSION NUMBER: 2001081341 PCTFULL ED 20020826 TITLE (ENGLISH): 9-OXA-EPOTHILON DERIVATIVES, METHOD FOR THE PRODUCTION AND USE THEREOF IN PHARMACEUTICAL PREPARATIONS TITLE (FRENCH): DERIVES DE 9-OXA-EPOTHILONE, LEUR PROCEDE DE PRODUCTION ET LEUR UTILISATION PHARMACEUTIQUE INVENTOR(S): SCHWEDE, Wolfgang; KLAR, Ulrich; SKUBALLA, Werner; BUCHMANN, Bernd; HOFFMANN, Jens; LICHTNER, Rosemarie PATENT ASSIGNEE(S): SCHERING AKTIENGESELLSCHAFT; SCHWEDE, Wolfgang; KLAR, Ulrich; SKUBALLA, Werner; BUCHMANN, Bernd; HOFFMANN, Jens: LICHTNER, Rosemarie DOCUMENT TYPE: Patient PATENT INFORMATION: NUMBER KIND DATE WO 2001081341 A2 20011101 DESIGNATED STATES W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF CG CI CM GA GN GW ML MR NE SN TD TG PRIORITY INFO.: DE 2000-100 20 899.1 20000420 APPLICATION INFO.: WO 2001-EP4551 A 20010419 ANSWER 4 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN ACCESSION NUMBER: 2001073103 PCTFULL ED 20020822 TITLE (ENGLISH): PREPARATION OF EPOTHILONE INTERMEDIATES TITLE (FRENCH): PREPARATION D'INTERMEDIAIRES D'EPOTHILONE INVENTOR(S): VITE, Gregory, D.; KIM, Soong-Hoon; HOeEFLE, Gerhard BRISTOL-MYERS SQUIBB COMPANY; PATENT ASSIGNEE(S): VITE, Gregory, D.; KIM, Soong-Hoon; HOeEFLE, Gerhard DOCUMENT TYPE: Patent PATENT INFORMATION:

DESIGNATED STATES ŢεJ •

NUMBER

AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG

KIND DATE WO 2001073103 A2 20011004

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                        TM TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ
                        SD SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH
                        CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ
                        CF CG CI CM GA GN GW ML MR NE SN TD TG
PRIORITY INFO.:
                        US 2000-60/191,975 20000324
                       WO 2001-US9620
                                            A 20010323
APPLICATION INFO.:
     ANSWER 5 OF 33
                        PCTFULL COPYRIGHT 2008 Univentio on STN
ACCESSION NUMBER:
                       2001070716 PCTFULL ED 20020822
TITLE (ENGLISH):
                       A PROCESS FOR THE PREPARATION OF EPOTHILONE
                        ANALOGS AND INTERMEDIATES
TITLE (FRENCH):
                        PREPARATION D'ANALOGUES ET D'INTERMEDIAIRES D'
                        EPOTHILONE
INVENTOR(S):
                        LI, Wen, Sen;
                        THORNTON, John, E.;
                        GUO, Zhenrong;
                        SWAMINATHAN, Shankar;
                        MCCONLOGUE, Gary, W.
PATENT ASSIGNEE(S):
                        BRISTOL-MYERS SQUIBB COMPANY;
                        LI, Wen, Sen;
                        THORNTON, John, E.;
                        GUO, Zhenrong;
                        SWAMINATHAN, Shankar;
                        MCCONLOGUE, Gary, W.
DOCUMENT TYPE:
                        Patent
PATENT INFORMATION:
                        NUMBER
                                          KIND DATE
                        WO 2001070716 A1 20010927
DESIGNATED STATES
                        AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR
                        CU CZ DE DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN
                        IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK
                        MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM
                        TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD
                        SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY
                        DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF
                        CG CI CM GA GN GW ML MR NE SN TD TG
PRIORITY INFO.: US 2000-09/528,526 20000320 APPLICATION INFO.: WO 2001-US7749 A 20010312
     (FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008)
     FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008
```

=> d his

W:

T. 1 2484 S EPOTHILON?

L2 159 S L1/AB OR L1/TI

L3 53 S L2 NOT PY>2001

L4 33 S COMBINATION AND L3

=> s 14 and (taxol or paclitaxel)

9622 TAXOL

272 TAXOLS

9705 TAXOL (TAXOL OR TAXOLS)

10390 PACLITAXEL

72 PACLITAXELS

10392 PACLITAXEL

(PACLITAXEL OR PACLITAXELS)

```
L5
          29 L4 AND (TAXOL OR PACLITAXEL)
=> s 15 and Her?
       988529 HER?
           29 L5 AND HER?
=> s 15 and (HER2 or HER-2)
          4722 HER2
        118696 HER
          1043 HERS
        119313 HER
                (HER OR HERS)
       1276185 2
         3260 HER-2
                (HER(W)2)
             1 L5 AND (HER2 OR HER-2)
=> d ibib abs
      ANSWER 1 OF 1
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ACCESSION NUMBER:
                       1999002514 PCTFULL ED 20020515
TITLE (ENGLISH):
                       EPOTHILONE DERIVATIVES
TITLE (FRENCH):
                       DERIVES D'EPOTHILONE
INVENTOR(S):
                       VITE, Gregory, D.:
                       BORZILLERI, Robert, M.;
                       KIM, Soong-Hoon;
                       JOHNSON, James, A.
                       BRISTOL-MYERS SOUIBB COMPANY
PATENT ASSIGNEE(S):
LANGUAGE OF PUBL.:
                       English
DOCUMENT TYPE:
                       Patent
PATENT INFORMATION:
                       NUMBER
                                         KIND DATE
                       WO 9902514
                                           A2 19990121
DESIGNATED STATES
      TaT •
                       AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE
                       ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC
                        LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU
                        SD SE SG SI SK SL TJ TM TR TT UA UG UZ VN YU ZW GH GM
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                       CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ
                       CF CG CI CM GA GN ML MR NE SN TD TG
PRIORITY INFO .:
                       US 1997-60/051,951
                                                19970708
                       US 1997-60/067,524
                                                19971204
APPLICATION INFO .:
                       WO 1998-US12550 A 19980616
      The present invention relates to compounds of formula (I), Q is selected
ABEN
       from the group
       consisting of (II), G is selected from the group consisting of alkyl,
       substituted alkyl, substituted
       or or unsubstituted aryl, heterocyclo, (III), W is O or NR15; X is O or
      H, H; Y is selected from the
       group consisting of O; H,OR16; OR17,OR17; NOR18; H,NOR19; H,NR20R21;
      H,H; or CHR22; OR170R17 can be
      a cyclic ketal; Z1 and Z2 are selected from the group consisting of CH2,
      O, NR23, S or SO2, wherein
      only one of Z and Z2 is a heteroatom; B1 and B2 are selected from the
      group consisting of OR24, or
      OCOR25, or 2CNR26R27; when B1 is H and Y is OH, H they can form a
      six-membered ring ketal or acetal;
      D is selected from the group consisting of NR28R29, NR30COR31 or
      saturated heterocycle R1, R2, R3,
      R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 and R27 are
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substituted alkyl, or aryl and when R1 and R2 are alkyl can be joined to
form a cycloalkyl; R3 and
R4 are alkyl can be joined to form a cycloalkyl; R9, R10, R16, R17, R24,
R25, and R31 are selected
from the group H, alkyl, or substituted alkyl; R8, R11, R12, R28, R30,
R32, R33, and R30 are
selected from the group consisting of H, alkyl, substituted alkyl, arvl,
substituted arvl.
cycloalkyl, or heterocyclo; R15, R23 and R29 are selected from the
group consisting of H, alkyl,
substituted alkyl, aryl, substituted aryl, cycloalkyl, heterocyclo,
R32C=O, R33SO2, hydroxy, O-alkyl
or O-substituted alkyl, the pharmaceutically acceptable salts thereof
and any hydrates, solvates or
geometric, optical and stereoisomers thereof, with the proviso that
compounds wherein: W and X are
both O; and R1, R2, R7 are H; and R3, R4, R6, are methyl; and R8, is H
or methyl; and Z1, and Z2,
are CH2; and G is 1-methyl-2-(substituted-4-thiazolyl)ethenyl; and Q is
as defined above are
excluded.
La presente invention concerne des composes de la formule (I) dans
laquelle O est selectionne
dans le groupe constitue par le groupement (II); G est selectionne dans
le groupe constitue par
alkyle, akyle substitue, aryle substitue ou insusbstitue, heterocyclo,
le groupement (III); W est O
ou NR15; X est O ou H,H; Y est selectionne dans le groupe constitue par
O; H, OR16; OR17, OR17; NOR18;
H, NOR19; H, NR20R21; H, H; ou CHR22; OR17, OR17 pouvant etre un cetal
cyclique; Z1 et Z2 sont
selectionnes dans le groupe constitue par CH2, O, NR23, S ou SO2, dans
lequel seuls Z et Z2 sont un
heteroatome; B1 et B2 sont selectionnes dans le groupe constitue par
OR24 ou OCOR25 ou O2CNR26R27;
et peuvent former ensemble un noyau cetal ou acetal a six chainons si B1
est H et Y est OH, H; D est
selectionne dans le groupe constitue par NR28R29, NR30COR31 ou un
heterocycle sature, R1, R2, R3,
R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 et R27 sont
selectionnes dans le groupe
constitue par H, alkyle, alkyle substitue ou aryle, et peuvent former
ensemble un cycloalkyle si R1
et R2 ou R3 et R4 sont alkyle; R9, R10, R16, R17, R24, R25 et R31 sont
selectionnes dans le groupe
constitue par H, alkyle ou alkyle substitue; R8, R11, R12, R28, R30,
R32, R33 et R30 sont
selectionnes dans le groupe constitue par H, alkyle, alkyle substitue,
aryle, aryle substitue,
cycloalkyle ou heterocyclo; R15, R23 et R29 sont selectionnes dans le
groupe constitue par H,
alkyle, alkyle substitue, arvle, arvle substitue, cycloalkyle ou
heterocyclo, R32C=O, R33SO2,
hydroxy, O-alkyle ou O-alkyle substitue, leurs sels pharmaceutiquement
acceptables ou leurs
eventuels hydrates, solvates ou isomeres geometriques, optiques, ou
stereoisomeres, a condition que
soient exclus les composes dans lesquels W et X sont tous deux O; et R1,
R2 et R7 sont H; et R3, R4
et R6 sont methyle; et R8 est H ou methyle; et Z1 et Z2 sont CH2; et G
est
```

selected from the group H, alkyl,

ABFR

1-methyl-2-(substitue-4-thiazolyl)ethenyle; et Q est tel que defini ci-dessus.

```
=> d his
     (FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008)
     FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008
           2484 S EPOTHILON?
L2
            159 S L1/AB OR L1/TI
L3
             53 S L2 NOT PY>2001
L4
             33 S COMBINATION AND L3
L5
             29 S L4 AND (TAXOL OR PACLITAXEL)
L6
            29 S L5 AND HER?
L7
             1 S L5 AND (HER2 OR HER-2)
=> s 16 and (HER2 or HER-2)
          4722 HER2
        118696 HER
          1043 HERS
        119313 HER
                (HER OR HERS)
       1276185 2
          3260 HER-2
                 (HER (W) 2)
1.8
             1 L6 AND (HER2 OR HER-2)
=> s 15 and (HER2 or HER-2)
          4722 HER2
        118696 HER
          1043 HERS
        119313 HER
                (HER OR HERS)
       1276185 2
          3260 HER-2
                (HER(W)2)
L9
             1 L5 AND (HER2 OR HER-2)
=> d ibib abs kwic
      ANSWER 1 OF 1
                        PCTFULL COPYRIGHT 2008 Univentio on STN
ACCESSION NUMBER:
                        1999002514 PCTFULL ED 20020515
                        EPOTHILONE DERIVATIVES
TITLE (ENGLISH):
TITLE (FRENCH):
                        DERIVES D'EPOTHILONE
INVENTOR(S):
                        VITE, Gregory, D.;
                        BORZILLERI, Robert, M.;
                        KIM, Soong-Hoon;
                        JOHNSON, James, A.
PATENT ASSIGNEE(S):
                        BRISTOL-MYERS SQUIBB COMPANY
LANGUAGE OF PUBL.:
                        English
DOCUMENT TYPE:
                        Patent
PATENT INFORMATION:
                                          KIND DATE
                        WO 9902514
                                            A2 19990121
DESIGNATED STATES
       w:
                        AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE
                        ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC
                        LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU
                        SD SE SG SI SK SL TJ TM TR TT UA UG UZ VN YU ZW GH GM
                        KE LS MW SD SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE
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CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ
                        CF CG CI CM GA GN ML MR NE SN TD TG
PRIORITY INFO.:
                       US 1997-60/051,951
                                                19970708
                       US 1997-60/067,524
                                                19971204
APPLICATION INFO.:
                       WO 1998-US12550
                                             A 19980616
      The present invention relates to compounds of formula (I), Q is selected
ABEN
       from the group
       consisting of (II), G is selected from the group consisting of alkyl,
       substituted alkyl, substituted
       or or unsubstituted arvl, heterocyclo, (III), W is O or NR15; X is O or
       H.H; Y is selected from the
       group consisting of O; H,OR16; OR17,OR17; NOR18; H,NOR19; H,NR20R21;
       H,H; or CHR22; OR170R17 can be
       a cyclic ketal; Z1 and Z2 are selected from the group consisting of CH2,
       O, NR23, S or SO2, wherein
       only one of Z and Z2 is a heteroatom; B1 and B2 are selected from the
       group consisting of OR24, or
       OCOR25, or 2CNR26R27; when B1 is H and Y is OH, H they can form a
       six-membered ring ketal or acetal;
       D is selected from the group consisting of NR28R29, NR30COR31 or
       saturated heterocycle R1, R2, R3,
       R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 and R27 are
       selected from the group H. alkvl.
       substituted alkyl, or aryl and when R1 and R2 are alkyl can be joined to
       form a cycloalkyl; R3 and
       R4 are alkyl can be joined to form a cycloalkyl; R9, R10, R16, R17, R24,
       R25, and R31 are selected
       from the group H, alkyl, or substituted alkyl; R8, R11, R12, R28, R30,
       R32, R33, and R30 are
       selected from the group consisting of H, alkyl, substituted alkyl, aryl,
       substituted aryl,
       cycloalkyl, or heterocyclo; R15, R23 and R29 are selected from the
      group consisting of H, alkyl,
       substituted alkyl, aryl, substituted aryl, cycloalkyl, heterocyclo,
      R32C=O, R33SO2, hydroxy, O-alkyl
      or O-substituted alkyl, the pharmaceutically acceptable salts thereof
       and any hydrates, solvates or
       geometric, optical and stereoisomers thereof, with the proviso that
       compounds wherein: W and X are
       both O; and R1, R2, R7 are H; and R3, R4, R6, are methyl; and R8, is H
      or methyl; and Z1, and Z2,
       are CH2; and G is 1-methy1-2-(substituted-4-thiazoly1)etheny1; and Q is
      as defined above are
       excluded.
       La presente invention concerne des composes de la formule (I) dans
```

## ABFR

laquelle Q est selectionne

dans le groupe constitue par le groupement (II); G est selectionne dans le groupe constitue par

alkyle, akyle substitue, aryle substitue ou insusbstitue, heterocyclo, le groupement (III); W est O

ou NR15; X est O ou H,H; Y est selectionne dans le groupe constitue par O; H, OR16; OR17, OR17; NOR18;

H, NOR19; H, NR20R21; H, H; ou CHR22; OR17, OR17 pouvant etre un cetal cyclique; Z1 et Z2 sont

selectionnes dans le groupe constitue par CH2, O, NR23, S ou SO2, dans lequel seuls Z et Z2 sont un

heteroatome; B1 et B2 sont selectionnes dans le groupe constitue par OR24 ou OCOR25 ou O2CNR26R27;

et peuvent former ensemble un noyau cetal ou acetal a six chainons si Bl est H et Y est OH, H; D est

selectionne dans le groupe constitue par NR28R29, NR30COR31 ou un heterocycle sature. R1, R2, R3,

```
R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 et R27 sont
selectionnes dans le groupe
constitue par H, alkyle, alkyle substitue ou aryle, et peuvent former
ensemble un cycloalkyle si R1
et R2 ou R3 et R4 sont alkyle; R9, R10, R16, R17, R24, R25 et R31 sont
selectionnes dans le groupe
constitue par H, alkyle ou alkyle substitue; R8, R11, R12, R28, R30,
R32, R33 et R30 sont
selectionnes dans le groupe constitue par H, alkyle, alkyle substitue,
arvle, arvle substitue,
cycloalkyle ou heterocyclo; R15, R23 et R29 sont selectionnes dans le
groupe constitue par H,
alkyle, alkyle substitue, aryle, aryle substitue, cycloalkyle ou
heterocyclo, R32C=0, R33S02,
hydroxy, O-alkyle ou O-alkyle substitue, leurs sels pharmaceutiquement
acceptables ou leurs
eventuels hydrates, solvates ou isomeres geometriques, optiques, ou
stereoisomeres, a condition que
soient exclus les composes dans lesquels W et X sont tous deux O; et R1,
R2 et R7 sont H; et R3, R4
et R6 sont methyle; et R8 est H ou methyle; et Z1 et Z2 sont CH2; et G
1-methyl-2-(substitue-4-thiazolyl)ethenyle; et 0 est tel que defini
ci-dessus.
EPOTHILONE DERIVATIVES
DERIVES D'EPOTHILONE
S Me
HOF
N3 ], ]'] '
0 Me
0 OH 0
I EpothiloneA R=H
II EpothiloneB R=Me
have been found to exert microtubule-stabilizing effects similar to
  TAXOL and hence cytotoxic activity against rapidly
proliferating cells,
such as, tumor cells or other hyperproliferative cellular disease, see
.Angew. Chem. Int. Ed. Engl.,. . .
The compounds of this invention, are also useful in combination
with known anti-cancer and cytotoxic agents and treatments, including
radiation. If formulated as a fixed dose, such combination
products
employ the compounds of this invention within the dosage range
described below and the other pharmaceutically active agent within its
approved dosage range. Compounds of formula V can be used
sequentially with known anticancer or cytotoxic agents and treatment,
including radiation when a combination formulation is
inappropriate.
Especially useful are cytotoxic drug combinations wherein the
second
drug chosen acts in a different phase of the cell cycle, e.g. S phase,
t.han
the present compounds of. . .
Synthase Inhibitors,
DNA Cross Linking Agents
Topoisomerase I and II Inhibitors
DNA Alkylating Agents
```

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TIFR

Ribonucleoside Reductase Inhibitors Cytotoxic Factors e.g. TNF-alpha or Growth factor inhibitors e.g. HER 2 receptor MAB's The present compounds may exist as multiple optical, geometric, and stereoisomers. Included within the present invention are all such isomers and. . . .

potency is

accomplished following a modified procedure of Swindell, et al., (see Swindell, C.S., Krauss, N.E., Horwitz, S.B., and Ringel, I. Biologically active taxol analogues with deleted A-ring side chain substituents and

variable C-2' configurations. J. Med. Chem. 34: 1176-1184, 1991). These modifications, in part, result. . .

cells were incubated at 37' form 72 hours at which time the tetrazolium dye, MTS at 333 gg/ml (final concentration), in combination

with the electron coupling agent phenazine methosulfate at 25 gm (final concentration) was added. A dehydrogenase enzyme in live cells

reduces the MTS. .

=>

---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

 COST IN U.S. DOLLARS
 SINCE FILE
 TOTAL

 FULL ESTIMATED COST
 23.30
 23.51

STN INTERNATIONAL LOGOFF AT 10:14:45 ON 07 JUL 2008